PATENT COOPERATION TREATY

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INTERNATIONAL PRELIMINARY REPORT ON PATENTABI

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

	olicant's or agent's fi 0/4-32804A	le reference	FOR FURTHER	ACTION	See Form PCT/IPEA/416				
	International application No. PCT/EP2004/003513 International filing date of the control of th			e (day/month/year)	Priority date (day/month/year) 04.04.2003				
A6 ⁻	International Patent Classification (IPC) or national classification and IPC A61K31/435, A61K31/436, A61K47/02, A61K47/10, A61K47/14, A61K47/44 Applicant								
NOVARTIS AG et al.									
1.	This report is th Authority under	e international prel Article 35 and tran	iminary examination is smitted to the applica	report, established by ant according to Article	this International Preliminary Examini	ing			
2.	This REPORT	consists of a total o	f 8 sheets, including	this cover sheet.					
3.	This report is al	so accompanied by	ANNEXES, compris	ing:					
	a. 🛭 sent to t	he applicant and to	the International Bur	eau) a total of 1 she	ets, as follows:				
	anu/	ets of the description for sheets containing inistrative Instruction	g recuircations author	rings which have been rized by this Authority	n amended and are the basis of this re (see Rule 70.16 and Section 607 of the	port he			
	Deyc	ets which supersed and the disclosure i plemental Box.	e earlier sheets, but v n the international ap	vhich this Authority co plication as filed, as i	onsiders contain an amendment that g ndicated in item 4 of Box No. I and the	oes ,			
	Sequenc	e nothing and/or table	es relateu thereto. In 1	indicate type and nun computer readable fo 02 of the Administrati	nber of electronic carrier(s)) , contain rm only, as indicated in the Suppleme ve Instructions).	ing a ntal			
4.	This report conta	ains indications rela	ating to the following i	tems:					
		Basis of the opini	on ·	•					
	☐ Box No. II ☐ Box No. III	Priority		•		1			
	☐ Box No. IV			ard to novelty, inventi	e step and industrial applicability				
	Box No. V Box No. V	Lack of unity of in Reasoned statem applicability: citation	nent under Article 35(2) with regard to nove supporting such state	Ity, inventive step or industrial				
	⊠ Box No. VI	Certain documen		capporaing odom otal					
	☐ Box No. VII	Certain defects in	the international app	lication					
	☐ Box No. VIII		ons on the internation						
Date	of submission of the	demand		Date of completion of	this report				
19.1	0.2004			14.11.2005					
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INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/003513

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	Bo	x No. I	Basis of t	ne repor	t							
1. With regard to the language, this report is based on the international application in th filed, unless otherwise indicated under this item.						the langu	lage in whic	h it wa				
		□ inte	rnational secondication of the	ige of a t arch (und e interna	slations from ranslation fur der Rules 12. Itional application	nished fo 3 and 23 ation (und	or the purp .1(b)) der Rule 12	oses of: 2.4)		languag	je ,	
2.	Hav	With regard to the elements* of the international application, this report is based on <i>(replacement sheets which</i> have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report):										
	Des	cription,	, Pages									
	1-13			as originally filed								
	Clai	ms, Nun	nbers									
	1-5				received on 1	9.10.2004	with letter	of 04.10.2	004			. •
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4.	Hau .	HOLDER	oort has bee n made, sind al Box (Rule	e they n	shed as if (so ave been cor	me of) th	e amendn to go beyo	nents ann and the dis	exed to thi	s report a s filed, as	and listed be indicated ir	elow 1 the
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INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2004/003513

		x No. III Non-establishment plicability	of o	pinion with regard to novelty, inventive step and industrial			
1.	The obv	The questions whether the claimed invention appears to be novel, to involve an inventive step (to be non-obvious), or to be industrially applicable have not been examined in respect of:					
		the entire international application,					
	Ø	claims Nos. 3					
		because:					
	\boxtimes	the said international application, or the said claims Nos. 3 relate to the following subject matter which does not require an international preliminary examination (specify):					
		see separate sheet					
		the description, claims or drawings (indicate particular elements below) or said claims Nos. are so unclear that no meaningful opinion could be formed (specify):					
		the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed.					
		no international search report has been established for the said claims Nos.					
İ		the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that:					
		the written form		has not been furnished			
				does not comply with the standard			
		the computer readable form		has not been furnished			
				does not comply with the standard			
I		the tables related to the nucleotide and/or amino acid sequence listing, if in computer readable form only, do not comply with the technical requirements provided for in Annex C-bis of the Administrative Instructions.					
	☐ See separate sheet for further details						

Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N)

Yes: Claims

No: Claims

1-5

Inventive step (IS)

Yes: Claims

No: Claims

1-5

Industrial applicability (IA)

Yes: Claims

1,2,4,5

No:

Claims

2. Citations and explanations (Rule 70.7):

see separate sheet

Box No. VI Certain documents cited

1. Certain published documents (Rule 70.10)

and/or

2. Non-written disclosures (Rule 70.9)

see separate sheet

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Re Item III

Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

Claim 3 relates to subject-matter considered by this Authority to be covered by the provisions of Rule 67.1(iv) PCT. Consequently, no opinion will be formulated with respect to the industrial applicability of the subject-matter of these claims (Article 34(4)(a)(I) PCT).

Re Item V

Reasoned statement under Rule 66.2(a)(ii) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

For the assessment of the present claim 3 on the question whether it is industrially applicable, no unified criteria exist in the PCT Contracting States. The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognize as industrially applicable the subject-matter of claims to the use of a compound in medical treatment, but may allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a medicament for a new medical treatment.

- 1 The following documents are referred to in this communication:
 - D1: WO 96/13249 A (SANDOZ AG; SCHMOOK FRITZ (AT); POPP XUE PING (CH); JACKMAN MARTIN (CH) 9 May 1996 (1996-05-09)
 - D2: WO 00/32234 A (NOVARTIS ERFIND VERWALT GMBH; NOVARTIS AG (CH); KRIWET KATRIN (DE); R) 8 June 2000 (2000-06-08)
 - D3: KAPP A ET AL: "LONG-TERM MANAGEMENT OF ATOPIC DERMATITIS IN INFANTS WITH TOPICAL PIMECROLIMUS, A NONSTEROID ANTI-INFLAMMATORY DRUG" JOURNAL OF ALLERGY AND CLINICAL IMMUNOLOGY, MOSBY YEARLY BOOK, INC, US, vol. 110, no. 2, August 2002 (2002-08), pages 277-284, XP009032310 ISSN: 0091-6749
 - D4: WO 97/25977 A (CIBA GEIGY AG; TIEMESSEN HARRY (DE)) 24 July 1997

(1997-07-24)

D5: EP-A-0 812 588 (YOSHITOMI PHARMACEUTICAL) 17 December 1997 (1997-12-17)

D6: EP-A-1 273 288 (NOVARTIS ERFIND VERWALT GMBH; NOVARTIS AG (CH)) 8 January 2003 (2003-01-08)

D7: GB-A-2 327 610 (NOVARTIS AG) 3 February 1999 (1999-02-03)

D8: EP-A-1 064 942 (FUJISAWA PHARMACEUTICAL CO) 3 January 2001 (2001-01-03)

D9: US 2002/044967 A1 (IBUKI RINTA ET AL) 18 April 2002 (2002-04-18)

D10: WO 2004/016289 A (NOVARTIS PHARMA GMBH; NOVARTIS AG (CH); SEKKAT NABILA (CH); KRIWET KA) 26 February 2004 (2004-02-26)

D11: WO 03/074054 A (NOVARTIS PHARMA GMBH; NOVARTIS AG (CH); BABIOLE SAUNIER MAGGY (FR); B) 12 September 2003 (2003-09-12)

The present application does not meet the criteria of Article 33(1) PCT, because the subject-matter of claims 1 to 5 is not new in the sense of Article 33(2) PCT.

WO-A-9613249 provides a topical composition, in the form of an emulsion, that comprises a compound of the FK506 class: 33-epi-chloro-33-desoxy-ascomycin; and glycerol; an unsaturated fatty alcohol and water and also a topical pharmaceutical composition that comprises a macrolide in suspension. In a further aspect, this invention provides the use of an unsaturated fatty alcohol to stabilise the 33-epi-chloro-33-desoxy-ascomycin in a pharmaceutical composition.

The topical compositions defined above are useful in the treatment of inflammatory and hyperproliferative skin diseases and of cutaneous manifestations of immunologically mediated diseases. Examples of such diseases are psoriasis, atopic dermatitis, contact dermatitis and further eczematous dermatitises, seborrhoeic dermatitis, Lichen planus. Pemphigus, bullous Pemphigoid, Epidermolysis bullosa, urticaria. angioedemas, vasculitides, erythemas. cutaneous eosinophilias. Lupus erythematous and Alopecia areata. See in particular the examples 23, 24, 25 and 26 and also claims 1 to 8 and page 5, first paragraph.

This teaching anticipates the subject-matter of claims 1 to 5.

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WO-A-0032234 relates to topical compositions comprising ascomycins, such as 33-epi-chloro-33-desoxy-ascomycin (see page 2) and a carrier vehicle such as cetyidimethicone copolyol, polyglyceryl-4-isostearate or glycerol (see the description page 11).

This teaching anticipates the subject-matter of claims 1 to 5.

WO-A-9725977 relates to a process for preparing an emulsion composition comprising 33-epi-chloro-33-desoxy-ascomycin or a derivative thereof as active agent and a stabiliser selected from a phospholipid, a glycolipid, a sphingolipid, a diacylphosphatidyl glycerol, an egg-phosphatidylglycerol, a soy-phosphatidyglycerol, a diacyl-phosphatidylglycerol, or a salt thereof; or a saturated, mono- or di-unsaturated (C12-24) fatty acid, or a salt thereof, and c) an organic solvent. The composition is used for injection and in the treatment of atopic dermatitis, contact dermatitis and further eczematous dermatitises, seborrhoeic dermatitis.

This teaching anticipates the subject-matter of claims 1 to 5.

EP-A-1 273288 discloses an emulsion composition comprising 33-epi-chloro-33-desoxy-ascomycin or derivative thereof and an emollient selected from a phospholipid, a glycolipid, a sphingolipid, a diacylphosphatidyl glycerol, an egg-phosphatidylglycerol, a soy-phosphatidylglycerol, a diacyl-phosphatidylglycerol, or a salt thereof. This teaching anticipates the subject-matter of claims 1 to 5.

D7 discloses a pharmaceutical composition for topical application comprising a macrolide, preferably 33-epi-chloro-33-desoxy-ascomycin which is stabilised by the presence of glycerol monostearate. The composition is used for the treatment of plaque psoriasis. This teaching anticipates the subject-matter of claims 1 to 5.

D8 relates to sustained release of macrolide compounds preferably 33-epi-chloro-33-desoxy-ascomycin. Macrolide compounds are used for preventing bone marrow transplant rejection or organ transplant rejection and for treating and preventing autoimmune disorders, inflammatory disorders (such as eczema, atopic dermatitis and

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (SEPARATE SHEET)

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myocarditis). These compositions are administered orally.

US2002044967 provides an oral formulation of a macrolide compound where the dissolution of the macrolide compound is under sustained release; and a sustained-release formulation containing a composition in solid solution, where the macrolide compound is present at an amorphous state in a solid base adsorbate of fk506, tacrolimus, ascomycin, ethyl rapamycin, desoxyascomycin, etc...

The present application does not meet the criteria of Article 33(1) PCT, because the subject-matter of claims 1 to 5 does not involve an inventive step in the sense of Article 33(3) PCT.

Re Item VI

Certain documents cited

The current assessment is based on the assumption that all claims enjoy priority rights from the filing date of the priority document. If it later turns out that this is not correct, the documents P,X cited in the international search report could become relevant to assess whether claims 1 to 5 satisfy the criteria set forth in Article 33(1) PCT.

WO2004016289 published 20040226

relates to single-phase topical liquid or semi-solid pharmaceutical compositions substantially free of ethanol and water and comprising an ascomycin, such as 33-epi-chloro-33-desoxy-ascomycin in a carrier vehicle.

WO03074054 published 20030912

relates to topical ophthalmic compositions comprising an ascomycin such as 33-epi-chloro-33-desoxy-ascomycin e.g. for the treatment of inflammatory diseases such as blepharitis.

Applicants have now found that ophthalmic compositions comprising an ascomycin and a carrier comprising a medium chain fatty acid triglyceride and/or isopropyl myristate are highly efficient and well tolerated by the ocular tissue.



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04725324 amended: 4-Oct-2004

Patent Claims

- 1. A pharmaceutical composition comprising 33-epichloro-33-desoxyascomycin in combination or association with an emollient selected from the group consisting of dimethicone, glycerol and isostearylstearate together with at least one pharmaceutically acceptable diluent or carrier.
- 2. A pharmaceutical composition of claim 1 wherein the emollient is present in an amount from about 10% to about 5000% w/w of the amount of 33-epichloro-33-desoxyascomycin.
- 3. A method of treatment of a dermatological or mucosal disease in a subject suffering from such a disease comprising co-administering synergistically effective amounts of a composition of claim 1.
- 4. A process for the preparation of a composition of any one of claims 1 or 2 comprising mixing 33-epichloro-33-desoxyascomycin and an emollient selected from the group consisting of dimethicone, glycerol and isostearylstearate in combination or association with at least one pharmaceutically acceptable diluent or carrier.
- 5. A kit of parts comprising 33-epichloro-33-desoxyascomycin and an emollient selected from the group consisting of dimethicone, glycerol and isostearylstearate in separate unit dosage forms, together with instructions for use.

IL/4-Oct-2004

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